AMENDMENTS TO THE CLAIMS

1. - 19. (cancelled).

 (currently amended) A compound represented by the formula (III), a salt thereof or a hydrate of them.

wherein

 R^1 designates a group represented by the formula -(CO)_h-(NR^a)_J-(CR^b=CR^c)_k-Ar (wherein R^a , R^b and R^c each independently designate a hydrogen atom, halogen atom, hydroxyl group, an optionally substituted C_{1-6} alkyl group, an optionally substituted C_{2-6} alkenyl group, an optionally substituted C_{1-6} alkylthio group, an optionally substituted C_{2-6} alkenylthio group, an optionally substituted C_{1-6} alkylthio group, an optionally substituted C_{2-6} alkenylthio group, an optionally substituted C_{3-8} cycloalkenyl group, an optionally substituted C_{4-1} aryl group or an optionally substituted C_{5-1} aryl group or an optionally substituted C_{6-14} aryl group

Rd, Re and Rf each independently designate a hydrogen atom, halogen atom, hydroxyl

group, cyano group, nitro group, carboxyl group, an optionally substituted C_{1-6} alkyl group, an optionally substituted C_{1-6} alkoxy group, an optionally substituted C_{2-7} acyl group, $-CO-NR^{2a}R^{2b}$, $-NR^{2b}CO-R^{2a}$ or $-NR^{2a}R^{2b}$ (wherein R^{2a} and R^{2b} each independently designate a hydrogen atom or an optionally substituted C_{1-6} alkyl group), provided that at least one of R^d , R^s and R^f is not a hydrogen atom;

L designates a single bond, an optionally substituted C₁₋₆ alkylene group, an optionally substituted C₂₋₆ alkynylene group;

X designates a single bond, or a group represented by -NR 7 -, -O-, -CO-, -S-, -SO-, -SO₂-, -CO-NR 8 -Z-, -C(O)O-, -NR 8 -CO-Z-, -NR 8 -C(O)O-, -NR 8 -S-, -NR 8 -SO-, -NR 8 -SO₂-Z-, -NR 9 -CO-NR 10 -, -NR 9 -CS-NR 10 -, -S(O)m-NR 11 -Z-, -C(=NR 12)-NR 13 -, -OC(O)-, -OC(O)-NR 14 - or -CH₂-NR 8 -COR 7 - (wherein R 7 , R 8 , R 9 , R 10 , R 11 , R 12 , R 13 and R 14 each independently designate a hydrogen atom, halogen atom, hydroxyl group, an optionally substituted C₁₋₆ alkyl group, an optionally substituted C₂₋₆ alkenyl group, an optionally substituted C₁₋₆ alkylthio group, an optionally substituted C₁₋₆ alkenylthio group, an optionally substituted C₁₋₆ alkylthio group, an optionally substituted C₃₋₈ cycloalkyl group, an optionally substituted C₃₋₈ cycloalkyl group, an optionally substituted C₄₋₁₄ aryl group or an optionally substituted S- to 14-membered heteroaryl group, Z designates a single bond or an optionally substituted C₁₋₆ alkylene group, and m designates 0, 1 or 2): and

Y designates any one group selected from the group consisting of a hydrogen atom, halogen atom, nitro group, hydroxyl group, cyano group, carboxyl group or an optionally

substituted C_{1-6} alkyl group, an optionally substituted C_{2-6} alkenyl group, an optionally substituted C_{2-6} alkenyl group, an optionally substituted C_{3-8} cycloalkenyl group, an optionally substituted C_{3-8} cycloalkenyl group, an optionally substituted C_{3-8} cycloalkenyl group, an optionally substituted C_{4-1} aryl group, an optionally substituted amino group and a group represented by the formula -W-R¹⁵(wherein W designates C_{4-1} aryl group, an optionally substituted C_{4-1} aryl group, an optionally substituted amino group, an optionally substituted C_{4-1} aryl group or an optionally substituted C_{4-

21. (cancelled).

- 22. (original) The compound according to claim 20, a salt threof or a hydrate of them, wherein either one of R^d, R^e and R^f is a halogen atom or an optionally substituted C₁₋₆ alkoxy group.
- 23. (currently amended) The compound according to $\frac{20 \text{ or claim } 20}{20 \text{ or claim } 22}$ any one of $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ any one of $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ any one of $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ any one of $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ and $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ any one of $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ any one of $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ any one of $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ any one of $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ any one of $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ any one of $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ any one of $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ any one of $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ any one of $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ any one of $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ any one of $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ any one of $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ any one of $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ any one of $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ any one of $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ any one of $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ and $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ and $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ and $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ and $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ and $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ and $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ and $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ and $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ and $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ and $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ and $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ and $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ and $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ and $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ and $\frac{20 \text{ or claim } 22}{20 \text{ or claim } 22}$ and $\frac{20$

substituted 4- to 14-membered non-aromatic heterocyclic group, an optionally substituted C_{6-14} aryl group or an optionally substituted 5- to 14-membered heteroaryl group is excluded.

24. - 48. (cancelled).

49. (previously presented) The compound according to claim 20, a salt thereof or a hydrate of them, wherein L and X are a single bond, Y is a 5- to 6-membered heteroaryl group, and Y is a group optionally substituted with 1 to 3 group(s) selected from Substituent group a2 described in claim 43.

50. (previously presented) A pharmaceutical composition comprising the compound according to claim 20, a salt thereof or a hydrate of them, and a pharmaceutically acceptable carrier.

51. (previously presented) A c-Jun amino-terminal kinase (JNKs) inhibitor comprising the compound according to claim 20, a salt thereof or a hydrate of them.

52. (previously presented) A c-Jun amino-terminal kinase 1 (JNK 1), c-Jun amino-terminal kinase 2 (JNK 2) and/or c-Jun amino-terminal kinase 3 (JNK 3) inhibitor, comprising the compound according to claim 20, a salt thereof or a hydrate of them.

- 53. (previously presented) An agent for treating or preventing immunological diseases, inflammatory diseases or metabolic disorders, which comprises the compound according to claim 20, a salt thereof or a hydrate of them.
- 54. (previously presented) An agent for treating or preventing neurodegenerative diseases, which comprises the compound according to claim 20, a salt thereof or a hydrate of them.
- 55. (previously presented) An agent for treating or preventing Alzheimer's disease, Parkinson's disease, Huntington's chorea, amyotrophic lateral sclerosis, multiple sclerosis or spinocerebellar degeneration, which comprises the compound according to claim 20, a salt thereof or a hydrate of them.
 - 56. 58. (cancelled).
- 59. (previously presented) A method for treating or preventing a disease based on JNK 3 action against which inhibition of a c-Jun amino-terminal kinase 3 (JNK 3) is effective for prevention or treatment, immunological diseases, inflammatory diseases, metablic disorders and/or neurodegenerative diseases, which comprises administering a pharmacologically effective amount of the compound according to claim 20, a salt thereof or a hydrate of them to a patient.

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60. (previously presented) A method for treating or preventing a disease based on JNK action against which inhibition of a c-Jun amino-terminal kinase (JNK) is effective for prevention or treatment, immunological diseases, inflammatory diseases, metablic disorders or neurodegenerative diseases, which comprises administering a pharmacologically effective amount of the compound according to claim 20, a salt thereof or a hydrate of them to a patient.

61. (currently amended) The method according to claim <u>60</u> [[20]], wherein the disease is Alzheimer's disease, Parkinson's disease, Huntington's chorea, amyotrophic lateral sclerosis, multiple sclerosis or spinocerebellar degeneration.

62. (new) A method for treating a disease based on JNK action against which inhibition of a c-Jun amino-terminal kinase (JNK) is effective, wherein said disease is Alzheimer's disease, Parkinson's disease, Huntington's chorea, amyotrophic lateral sclerosis, multiple sclerosis, or spinocerebellar degeneration, which method comprises administering a pharmacologically effective amount of the compound according to claim 20, a salt thereof or a hydrate of them to a patient.